EXACT-RX SODIUM SULFACETAMIDE WASH 10% - sodium sulfacetamide liquid Exact-Rx, Inc.

Disclaimer: This drug has not been found by FDA to be safe and effective, and this labeling has not been approved by FDA. For further information about unapproved drugs, click here.

Exact-Rx Sodium Sulfacetamide Wash 10%

INDICATIONS: Sodium Sulfacetemide 10% Wash is intended for topical application in the following scaling dermatoses: seborrheic dermatisis and seborrhea sicca (dandruff). It also is indicated for the treatment of secondary bacterial infections of the skin due to organisms susceptible to sulfonamides.

DIRECTIONS FOR USE: Wash affected areas twice daily (morning and evening) or as directed by your physician. Rinse thoroughly and pat dry. **See package insert for complete product information**.

FOR EXTERNAL USE ONLY. NOT FOR INTRAVAGINAL OR OPHTHALMIC USE. (KEEP AWAY FROM EYES).

KEEP THIS AND ALL MEDICATION OUT OF REACH OF CHILDREN.

In case of accidental ingestion contact a poison control center immediately. Keep container tightly closed.

CONTRAINDICATIONS: Sodium Sulfacetamide 10% Wash is

contraindicated in persons with know or suspected hypersensitivity to sulfonamides.

Each gram contains 100 mg of sodium sulfacetamide USP in a

vehicle consisting of: citric acid, cocamidopropyl betaine, disodium EDTA,

glyceryl stearate, methylparaben, PEG-6 caprylic/capric

glycerides, PEG-60 almond glycerides, PEG-150 pentaerythrityl

tetrastearate, polysorbate 60, purified water, sodium lauryl

sulfate, sodium thio sulfate and xanthan gum.

Store at 25C (77F); excursions permitted to 15 to 30C (59 to

 $86F)\!.$ See USP Controlled Room. Protect from freezing.

See bottle for lot number and expiration date

Manufactured in the U.S.A. for

Exact-Rx, Inc., Melville, NY 11747

SODIUM SULFACETAMIDE 10% WASH

(sodium sulfacetamide 10%)

Rx Only

FOR EXTERNAL USE ONLY.

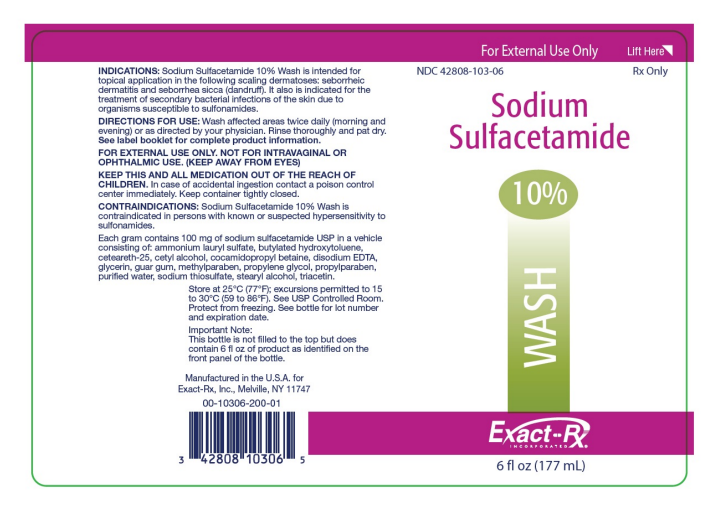
NOT FOR OPHTHALMIC USE.

Description: Each gram contains 100 mg of sodium sulfacetamide USP in a vehicle consisting of: ammonium lauryl sulfate, butylated hydroxytoluene, ceteareth-25, cetyl alcohol, cocamidopropyl betaine, disodium EDTA, glycerin, guar gum, methylparaben, propylene glycol, propylparaben, purified water, sodium thiosulfate, stearyl alcohol, triacetin.

HOW SUPPLIED: Sodium Sulfacetamide Wash 10% is available in a 6 fl oz (170 mL) bottle, NDC 42808-101-06, and in a 12 fl oz (354.8 mL) bottle, NDC 42808-101-12.

Manufactured in the U.S.A. for Exact-Rx, Inc., Melville, NY 11747

00-101-205-00 Iss:12/16



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DIRECTIONS FOR USE: Wash affected areas twice daily (morning and evening) or as directed by your physician. Rinse thoroughly and pat dry. See label booklet for complete product information.

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CONTRAINDICATIONS: Sodium Sulfacetamide 10% Wash is contraindicated in persons with known or suspected hypersensitivity to sulfonamides.

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Store at 25°C (77°F); excursions permitted to 15 to 30°C (59 to 86°F). See USP Controlled Room. Protect from freezing. See bottle for lot number and expiration date.

Important Note:

This bottle is not filled to the top but does contain 12 fl oz of product as identified on the front panel of the bottle.

Manufactured in the U.S.A. for Exact-Rx, Inc., Melville, NY 11747 00-10312-200-01



NDC 42808-103-12

Rx Only

Sodium Sulfacetamide



EXACT-RX SODIUM SULFACETAMIDE WASH 10%

sodium sulfacetamide liquid

Product Information			
Product Type	HUMAN PRESCRIPTION DRUG	Item Code (Source)	NDC:42808-103
Route of Administration	TOPICAL		

Active Ingredient/Active Moiety			
Ingredient Name	Basis of Strength	Strength	
SULFACETAMIDE SO DIUM (UNII: 4NRT660 KJQ) (SULFACETAMIDE -	SULFACETAMIDE	100 mg	
UNII:4965G3J0F5)	SODIUM	in 1 mL	

Inactive Ingredients			
Ingredient Name	Strength		
AMMO NIUM LAURYL SULFATE (UNII: Q7AO2R1M0B)			
BUTYLATED HYDRO XYTO LUENE (UNII: 1P9 D0 Z171K)			
COCAMIDO PRO PYL BETAINE (UNII: 50 CF30 11KX)			
CETEARETH-25 (UNII: 8FA93U5T67)			
CETYL ALCOHOL (UNII: 936JST6JCN)			
EDETATE DISO DIUM ANHYDRO US (UNII: 8 NLQ36 F6 MM)			
GLYCERIN (UNII: PDC6A3C0OX)			
GUAR GUM (UNII: E8911637KE)			
METHYLPARABEN (UNII: A2I8 C7HI9 T)			
PROPYLENE GLYCOL (UNII: 6 DC9 Q167V3)			
PROPYLPARABEN (UNII: Z8 IX2SC1OH)			
WATER (UNII: 059QF0KO0R)			
SODIUM THIO SULFATE (UNII: HX1032V43M)			
STEARYL ALCOHOL (UNII: 2KR89I4H1Y)			
TRIACETIN (UNII: XHX3C3X673)			

]	Packaging					
#	# Item Code	Package Description	Marketing Start Date	Marketing End Date		
1	NDC:42808-103- 06	177 mL in 1 BOTTLE; Type 0: Not a Combination Product	08/01/2011			
2	NDC:42808-103-12	354.8 mL in 1 BOTTLE; Type 0: Not a Combination Product	08/01/2011			

Marketing Information				
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date	
unapproved drug other		08/01/2011		

Labeler - Exact-Rx, Inc. (137953498)

Revised: 5/2020 Exact-Rx, Inc.